

Oges User Guide, Version 2.0, A Solver for Steady State Boundary Value Problems on Overlapping Grids

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Abstract: We describe the Overture class Oges, an “Overlapping Grid Equation Solver”, that can be used for the solution of sparse matrix equations on overlapping grids such that those created by the grid generator Ogen. Oges acts as a front end to a variety of sparse matrix solvers including direct sparse solvers such as those from Yale or Harwell or iterative solvers (from SLAP and PETSc) that use algorithms such as conjugate gradient or GMRES.

To use Oges one must first generate a system of equations (usually defining a PDE boundary value problem) using the ‘coefficient matrix’ grid functions and the Overture operator classes. Oges will take a coefficient matrix generated in this way and then call the appropriate sparse matrix solver. Oges can be easily extended to use a new Sparse matrix package.

This document is available from the **Overture** home page, <http://www.llnl.gov/casc/Overture>.

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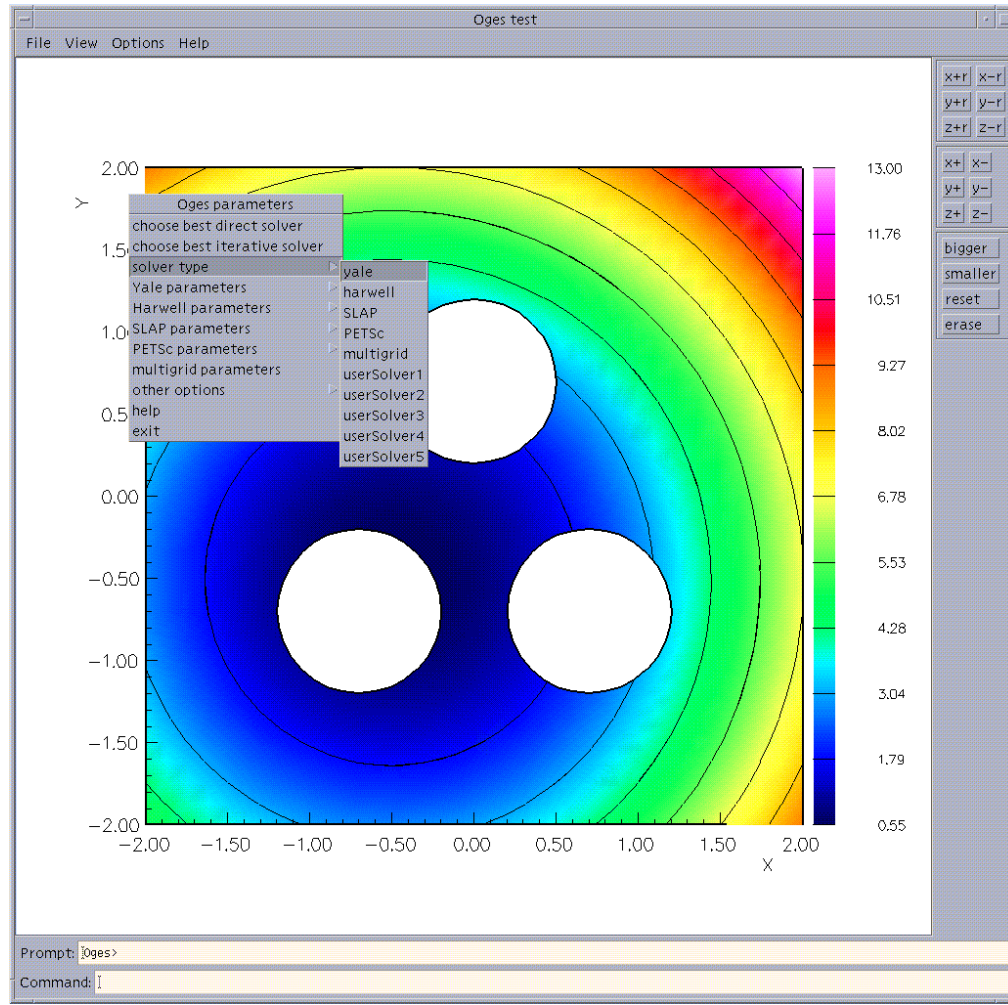


Figure 1: The ‘to ges’ program can be used to test Oges

1 Introduction

We describe the Overture class Oges, an “Overlapping Grid Equation Solver”, that can be used for the solution of sparse matrix equations on overlapping grids such that those created by the grid generator Ogen. Oges acts as a front end to a variety of sparse matrix solvers. Currently we have support for

Yale : direct sparse matrix package (no pivoting).

Harwell : direct sparse matrix package with partial pivoting.

SLAP : The Sparse Linear Algebra Package from Greenbaum and Seager, an iterative solver package, includes conjugate gradient and gmres solvers.

PETSc : The Portable Extensible Toolkit for Scientific computations[1] from iterative solver package, includes conjugate gradient and gmres solvers in addition to many others.

By changing one or two parameters the user may easily try a different solver. For example, although Yale is in general faster than Harwell, the latter, which does pivoting, may be better for some problems. The SLAP and PETSc iterative solvers may be especially useful for very large problems when storage is at a premium.

Oges can be easily extended by you to use a new Sparse matrix package.

To use Oges one must first generate a system of equations (usually defining a PDE boundary value problem) using the ‘coefficient matrix’ grid functions and the Overture operator classes [4][3][2]. A ‘coefficient matrix’ is stored in a `realCompositeGridFunction`. Typically the creation of a PDE boundary value problem will look something like

```

CompositeGrid cg(...);
realCompositeGridFunction coeff(...);
CompositeGridOperators op(cg);
...
coeff=op.laplacianCoefficients();           // form the laplace operator

coeff.applyBoundaryConditionCoefficients(0,0,dirichlet, allBoundaries);
coeff.applyBoundaryConditionCoefficients(0,0,extrapolate,allBoundaries);
coeff.finishBoundaryConditions();

```

Oges will take a coefficient matrix generated in this way and then call the appropriate sparse matrix solver. Usually this will involve converting the ‘coefficient matrix’ representation to some other representation such as a compressed-row storage format (this is done automatically by Oges).

Given a coefficient matrix, Oges can be used as follows

```

Oges solver(cg);                      // build a solver
solver.set(OgesParameters::THEsolverType,OgesParameters::yale); // use the yale solver
..or..
solver.set(OgesParameters::THEsolverType,OgesParameters::PETSc); // ...or.. use PETSc

solver.setCoefficientArray( coeff ); // supply coefficients
realCompositeGridFunction u(cg),f(cg); // build solution and right-hand-side

... assign f ...

u=0.;                                // initial guess for iterative solvers
solver.solve(u,f);                   // solve the equations
...

```

Generally one must also set other parameters such as the convergence tolerance, preconditioner, etc, when using iterative solvers such as SLAP or PETSC.

The global variable `Oges::debug` is a bit flag that generates various diagnostic output from Oges. Setting `Oges::debug=63` ($63=1+2+4+8+16+32$) will generate lots of debugging output. Setting `Oges::debug=3` will generate only some debugging output.

2 Example Codes

The file `Overture/tests/toges.C` is a test program for Oges. See the files `Overture/tests/tcm.C`, `tcm2.C`, `tcm3.C`, `tcm4.C` for examples of working codes. See also the examples presented in the Overture primer[5]. The flow solver `OverBlown` [6] also uses Oges. One could look at the source files for `OverBlown` for further examples.

3 Oges Parameters

Solver dependent parameters are found in the `OgesParameters` class. It is a container class for such parameters are the type of solver, type of preconditioner, convergence tolerance etc. `Oges` contains an `OgesParameters` object to hold these parameters. Parameters can be set by directly using the `Oges` `set` functions. This will indirectly set the values in an `OgesParameters` object contained in an `Oges` object. Alternatively one can first create an `OgesParameters` object, set parameters in that object and then provide the `OgesParameters` object to `Oges` using the `setParameters` function (which will copy you values into it's local version). Parameters can also be set interactively by calling the `Oges` `update` function or the `OgesParameters` `update` function.

3.1 operator=

```
OgesParameters&
operator=(const OgesParameters& x)
```

Description: deep copy of data.

3.2 getSolverName

```
aString
getSolverName() const
```

Description: Return the name of the solver, a composite of the solver type, method and preconditioner.

3.3 getSolverTypeName

```
aString
getSolverTypeName(SolverEnum solverType = defaultSolver) const
```

Description: Return the name of the solverType such as "yale", "harwell", "SLAP", ... By default return the name of the currently chosen solver.

solverType (input) : return the name of this solver type. By default return the name of the currently chosen solver.

3.4 getSolverMethodName

```
aString
getSolverMethodName(SolverMethodEnum solverMethodType = defaultSolverMethod) const
```

Description: Return the name of the solver method such as "gmres". By default return the name of the currently chosen method.

solverMethodType (input):

3.5 getPreconditionerName

```
aString
getPreconditionerName(PreconditionerEnum preconditionerType = defaultPreconditioner) const
```

Description: Return the name of the preconditioner. By default return the name of the currently chosen preconditioner.

preconditionerType (input):

3.6 getMatrixOrderingName

```
aString
getMatrixOrderingName(MatrixOrderingEnum matrixOrderingType = defaultMatrixOrdering) const
```

Description: Return the name of the matrix ordering. By default return the name of the currently chosen matrix ordering.

matrixOrderingType (input) :

3.7 set(OptionEnum , int)

int

set(OptionEnum option, int value = 0)

Description: Set an int option from the OptionEnum.

```

enum OptionEnum
{
    THEabsoluteTolerance,
    THEbestIterativeSolver, // choose the 'best' iterative solver and options.
    THEbestDirectSolver,   // choose the 'best' direct solver and options.
    THEcompatibilityConstraint,
    THEfillinRatio,
    THEfillinRatio2,
    THEfixupRightHandSide,
    THEgmresRestartLength,
    THEharwellPivotingTolerance,
    THEincompleteLUExpectedFill,
    THEiterativeImprovement,
    THEkeepCoefficientGridFunction, // keep a reference to the user's coeff grid function
    THEkeepSparseMatrix,          // keep ia,ja,a sparse matrix even it not needed by
    THEmatrixCutoff,
    THEmatrixOrdering,
    THEmaximumInterpolationWidth,
    THEmaximumNumberOfIterations,
    THEminimumNumberOfIterations,
    THEnullVectorScaling,
    THEnumberOfIncompleteLULevels,
    THEsolveForTranspose,
    THEpreconditioner,
    THEremoveSolutionAndRHSVector, // de-allocate sol and rhs vector after every solve
    THEremoveSparseMatrixFactorization, // de-allocate any factorization info after every solve.
    THErelativeTolerance,
    THErescaleRowNorms,
    THEsolverType,
    THEsolverMethod,
    THEtolerance,
    THEzeroRatio
};

```

3.8 set(OptionEnum , float)

int

set(OptionEnum option, float value)

Description: Set a real valued option from the OptionEnum.

3.9 set(OptionEnum , double)

int

set(OptionEnum option, double value)

Description: Set a real valued option from the OptionEnum.

3.10 set(SolverEnum)

```
int  
set( SolverEnum option )
```

Description: Set the solver, a value from the SolverEnum.

```
enum SolverEnum  
{  
    defaultSolver,  
    sor,  
    yale,  
    harwell,  
    SLAP,  
    PETSc,  
    userSolver1, // these are reserved for new user defined solvers.  
    userSolver2,  
    userSolver3,  
    userSolver4,  
    userSolver5  
};
```

3.11 set(SolverMethodEnum)

```
int  
set( SolverMethodEnum option )
```

Description: Set the solver method, a value from the SolverMethodEnum.

```
enum SolverMethodEnum  
{  
    richardson,  
    chebychev,  
    conjugateGradient,  
    cg=conjugateGradient, // cg= short PETSc name  
    biConjugateGradient,  
    bicg=biConjugateGradient,  
    conjugateGradientSquared,  
    cgs=conjugateGradientSquared,  
    biConjugateGradientSquared,  
    biConjugateGradientStabilized,  
    bcgs=biConjugateGradientStabilized,  
    generalizedMinimalResidual,  
    gmres=generalizedMinimalResidual,  
    transposeFreeQuasiMinimalResidual,  
    tfqmr=transposeFreeQuasiMinimalResidual,  
    transposeFreeQuasiMinimalResidual2, // tcqmr Tony Chan's version  
    tcqmr=transposeFreeQuasiMinimalResidual,  
    conjugateResidual,  
    cr=conjugateResidual,  
    leastSquares,  
    lsqr=leastSquares,  
    preonly,  
};
```

3.12 set(PreconditionerEnum)

int

set(PreconditionerEnum option)

Description: Set the preconditioner, a value from the PreconditionerEnum.

```
enum PreconditionerEnum
{
    noPreconditioner,
    jacobiPreconditioner,
    sorPreconditioner,
    luPreconditioner,
    shellPreconditioner,
    blockJacobiPreconditioner,
    multigridPreconditioner,
    eisenstatPreconditioner,
    incompleteCholeskyPreconditioner,
    incompleteLUPreconditioner,
    additiveSchwarzPreconditioner,
    slesPreconditioner,
    compositePreconditioner,
    redundantPreconditioner,
    diagonalPreconditioner,
    ssorPreconditioner
};
```

3.13 set(MatrixOrderingEnum)

int

set(MatrixOrderingEnum option)

Description: Set the matrix ordering, a value from the MatrixOrderingEnum.

```
enum MatrixOrderingEnum
{
    naturalOrdering,
    nestedDisectionOrdering,
    oneWayDisectionOrdering,
    reverseCuthillMcKeeOrdering,
    quotientMinimumDegreeOrdering,
    rowlengthOrdering
};
```

3.14 get(OptionEnum , int &)

int

get(OptionEnum option, int & value) const

Description: Get the value of an ‘int’ valued option.

3.15 get(OptionEnum , real &)

int

get(OptionEnum option, real & value) const

Description: Get the value of an ‘real’ valued option.

3.16 getOgmgParameters

OgmgParameters*
getOgmgParameters() const

Description: Return a pointer to the OgmgParameters object. This pointer may be NULL.

3.17 get from a data base

int
get(const GenericDataBase & dir, const aString & name)

Description: Get a copy of the OgesParameters from a database file

dir (input): get from this directory of the database.

name (input): the name of Oges on the database.

3.18 put to a data base

int
put(GenericDataBase & dir, const aString & name) const

Description: Output an image of OgesParameters to a data base.

dir (input): put onto this directory of the database.

name (input): the name of Oges on the database.

3.19 display

int
display(FILE *file = stdout)

Description: Print out current values of parameters

file (input) : print to this file (standard output by default).

3.20 update

int
update(GenericGraphicsInterface & gi, CompositeGrid & cGrid)

Description: Update parameters interactively.

gi: use this graphics interface.

cg: parameters will apply to this grid.

3.21 isAvailable(SolverEnum)

int
isAvailable(SolverEnum solverType)

Description: Return TRUE if a given solver (esp. PETSc) is available.

3.22 isSolverIterative

bool
isSolverIterative() const

Description: Return TRUE if the solver chosen is an iterative method

3.23 buildEquationSolvers

```
int  
buildEquationSolvers( SolverEnum solver)
```

Description: This function will build an equation solver of a particular type. This function is found in the Oges/buildEquationSolvers file. It is this file that you may have to copy and edit in order to turn on the availability solvers that are not distributed with Overture (such as PETSc).

4 Convergence criteria

There are many ways to define convergence criteria for iterative methods. The trick for Oges is to have a reasonable uniform way of defining a convergence tolerance for the different methods.

The standard PETSc convergence test is

$$\|r_k\|_2 < \max(rtol * \|r_0\|_2 <, atol) \quad \text{PETSc}$$

where

$$\|x\|_2 = \sqrt{\sum_i x_i^2}$$

The SLAP convergence test is somewhat different:

```
C ***** SLAP *****  
C *Description:  
C     SGMRES solves a linear system A*X = B rewritten in the form:  
C  
C     (SB*A*(M-inverse)*(SX-inverse))*(SX*M*X) = SB*B,  
C  
C     with right preconditioning, or  
C  
C     (SB*(M-inverse)*A*(SX-inverse))*(SX*X) = SB*(M-inverse)*B,  
C  
C     with left preconditioning, where A is an N-by-N real matrix,  
C     X and B are N-vectors, SB and SX are diagonal scaling  
C     matrices, and M is a preconditioning matrix. It uses  
C     preconditioned Krylov subspace methods based on the  
C     generalized minimum residual method (GMRES). This routine  
C     optionally performs either the full orthogonalization  
C     version of the GMRES algorithm or an incomplete variant of  
C     it. Both versions use restarting of the linear iteration by  
C     default, although the user can disable this feature.  
C  
C     The GMRES algorithm generates a sequence of approximations  
C     X(L) to the true solution of the above linear system. The  
C     convergence criteria for stopping the iteration is based on  
C     the size of the scaled norm of the residual R(L) = B -  
C     A*X(L). The actual stopping test is either:  
C  
C         norm(SB*(B-A*X(L))) .le. TOL*norm(SB*B),  
C  
C     for right preconditioning, or  
C  
C         norm(SB*(M-inverse)*(B-A*X(L))) .le.  
C             TOL*norm(SB*(M-inverse)*B),  
C  
C     for left preconditioning, where norm() denotes the euclidean  
C     norm, and TOL is a positive scalar less than one input by
```

C the user. If TOL equals zero when SGMRES is called, then a
C default value of 500*(the smallest positive magnitude,
C machine epsilon) is used. If the scaling arrays SB and SX
C are used, then ideally they should be chosen so that the
C vectors SX*X(or SX*M*X) and SB*B have all their components
C approximately equal to one in magnitude. If one wants to
C use the same scaling in X and B, then SB and SX can be the
C same array in the calling program.
C

5 Linking to PETSc

An example of linking to the PETSc libraries can be found in the `Overture/tests` directory. Type `'make tcm3p'` to build the `tcm3.C` test code with PETSc. Type `'tcm3p cic.hdf -solver=petsc'` to run the example on the grid `cic.hdf` with PETSc. This example assumes that the `PETSC_DIR`, `PETSC_ARCH` and `PETSC_LIB` environmental variables have been defined per instructions with the PETSc installation.

Here is an explanation of the steps required to build an Overture application with PETSc (as implemented in the above example). By default, the Overture library is unaware whether PETSc solvers are available. To use PETSc you should

1. Build or locate a version of PETSc. I have only built and linked Overture to the non-parallel version of PETSc. Link to the PETSc libraries (and `lapack`). I link to

```
petscLib = -L$(PETSC_LIB) -lpetscsles -lpetscdm -lpetscmat -lpetscvec -lpetsc \
-L/usr/local/lib -llapack -L$(PETSC_LIB) -lmpiuni
```

where `$(PETSC_LIB)` is the location of the PETSc libraries.

2. Copy the files `Oges/buildEquationSolvers.C` and `Oges/PETScEquationSolver.C` to your application directory and compile this file with the flags `-DOVERTURE_USE_PETSC` (or edit the file and define this variable inside with `#define OVERTURE_USE_PETSC`).
3. Link these new files, `buildEquationSolvers.o` and `PETScEquationSolver.o` with your application (ahead of the Overture library so that you get the new version) along with the PETSc libraries.

6 Adding a new sparse matrix solver to Oges

If you want to add a new sparse matrix solver to Oges you should look at one of the existing solvers, `YaleEquationSolver`, `HarwellEquationSolver`, `SlapEquationSolver` or `PETScEquationSolver`. These classes all derive from the base class `EquationSolver`. Oges contains a list of pointers to these `EquationSolver`'s. You will be able to add a new solver to this list. It will be known as `OgesParameters::userSolver1`, or `OgesParameters::userSolver2` etc., depending on how many new solvers have been added.

You should

1. Derive a new class from `EquationSolver`, copying one of the existing solvers (which ever is closest) to your new solver. Hopefully you can reuse parameters that already exist in `OgesParameters`.
2. Change the `Oges/buildEquationSolvers.C` file to ‘new’ the solver to have defined and add it to the list of `EquationSolver`'s. Change the other functions in `Oges/buildEquationSolvers.C` as appropriate.
3. Compile your files and the new version of `buildEquationSolvers.C` and link to these ahead of the Overture library when you build an executable.

7 Some More Details about Oges

In general, Oges expects that the user wants to solve one or more equations at each valid grid point on an overlapping grid. The number of equations that are given at each grid point is called the `numberOfComponents`. In the simple case only one equation, such as a discrete Laplace operator, is specified at each grid point, `numberOfComponents=1`. In a more complicated case there will be a system of equations at each grid point. For example, one may want to solve the biharmonic equation as a system of two Possion equations in which case `numberOfComponents=2`.

Oges will create a large sparse matrix where each unknown for the sparse matrix will correspond to a particular component `n`, at a particular grid point (`i1, i2, i3`) on a particular component grid, `grid`. Thus there is a mapping from `(n, i1, i2, i3, grid)` to a unique equation number. The member functions

```
int equationNo(int n, int i1, int i2, int i3, int grid)
intArray equationNo(int n, Index & I1, Index & I2, Index & I3, int grid)
```

give the equation number(s) for each grid point. For now the function `equationNo` is defined to use all the grid points in a given order. In the future, a user should be able to define this function in a different way. There is also a member function

```
void equationToIndex(int eqnNo, int n, int i1, int i2, int i3, int grid)
```

that maps an equation number, `eqnNo`, back to a grid point and component, `(n, i1, i2, i3, grid)` (i.e. it is the inverse of `equationNo`).

Sometimes extra unknowns and extra equations are required in order to specify a problem. For example, an eigenvalue problem has an extra unknown, the eigenvalue. An extra unknown may be added to the singular Neumann problem in order to create a nonsingular system. Extra unknowns are associated with grid points that are not used. The number of extra equations is specified with `setNumberOfExtraEquations`. Oges will find unused points that can be used for extra equations; the equation numbers for these points will be saved in `extraEquationNumber(i)`.

8 Oges Function Descriptions

8.1 default constructor

`Oges()`

Description: Default constructor.

8.2 setGridsToUse

```
int  
setGridsToUse( const IntegerArray & gridsToUse_ )
```

Description: Only solve the equations on some grids.

8.3 getMaximumResidual

```
real  
getMaximumResidual() const
```

Description: Return the maximum residual from the last solve.

8.4 get

```
int  
get( const GenericDataBase & dir, const aString & name)
```

Description: Get a copy of Oges from a database file

dir (input): get from this directory of the database.

name (input): the name of Oges on the database.

8.5 put

```
int  
put( GenericDataBase & dir, const aString & name) const
```

Description: Output an image of Oges to a data base.

dir (input): put onto this directory of the database.

name (input): the name of Oges on the database.

8.6 writeMatrixToFile

```
int  
writeMatrixToFile( aString filename )
```

Description: // Write the current solver matrix to the file `filename`. // The file consists of triplets $i, j, A(i, j)$ (without commas) // for each non-zero element of the matrix. // (Here i =row, j =column, and $A(i, j) = A_{ij}$ element of the matrix.)

Author: pf

8.7 writeMatrixGridInformationToFile

```
int  
writeMatrixGridInformationToFile( aString filename )
```

Description: Write the grid information about the current solver matrix to the file `filename`.

For each equation in the matrix, a line
is saved in the file with the following format:

```
ieq    grid    simpleClassify    fullClassify
```

where:

```
ieq=          equation number in the linear system  
grid=         grid number for this point
```

(In the classify flags, any non-negative value indicates a used
point. Negative values are equations with zero for the rhs)

```
simpleClassify=  
              -1=connecting grids (=interpolation,  
                           extrapolation, or periodic bdry)  
              0=hole point (unused)  
              1=interior (=discretization) point  
              2=boundary point  
                  (=boundary, ghostline, periodic)
```

```
fullClassify=  
              interior=      1,  
              boundary=      2,  
              ghost1=        3,  
              ghost2=        4,  
              ghost3=        5,  
              ghost4=        6,  
              interpolation= -1,  
              periodic=       2,  
              extrapolation= -3,  
              unused=         0
```

Author: pf

8.8 writePetscMatrixToFile

```
int  
writePetscMatrixToFile( aString filename,  
                       realCompositeGridFunction & u,  
                       realCompositeGridFunction & f)
```

Description: Only available when linked with PETSc (-DOVERTURE_USE_PETSC)

Write the current solver matrix to the file `filename`. Uses the PETSc binary format. Supply `u,f` as to 'solver', the RHS corresponding to `f` is also saved in the matrix file.

Author: pf

8.9 canSolveInPlace

```
bool
canSolveInPlace() const
```

Description: Return TRUE if the rhs and sol vectors can be the same.

8.10 setCoefficientArray

```
int
setCoefficientArray( realCompositeGridFunction & coeff0 )
```

Purpose: Supply a coefficient grid function to be used to discretize the equations.

coeff0 (input): Here are the coefficients. Oges will keep a reference to this grid function.

8.11 setCoefficientArray

```
int
setCoefficientArray( realMappedGridFunction & coeff0 )
```

Purpose: Supply a coefficient grid function (single grid only) to be used to discretize the equations.

coeff0 (input): Here are the coefficients. Oges will keep a reference to this grid function.

8.12 setEvaluateJacobian

```
void
setEvaluateJacobian( const int evaluateJacobian0 )
```

Purpose: ?

8.13 setGrid

```
void
setGrid( CompositeGrid & cg0 )
```

Purpose: Supply a CompositeGrid to Oges. Use this routine, for example, if an Oges object was created with the default constructor. Call this routine before calling initialize.

cg0 (input): Oges will keep a reference to this grid.

8.14 setGrid

```
void
setGrid( MappedGrid & mg )
```

Purpose: Supply a MappedGrid to Oges. Use this routine, for example, if an Oges object was created with the default constructor. Call this routine before calling initialize.

mg (input): Oges will keep a reference to this grid.

8.15 set(OptionEnum,int)

```
int
set( OptionEnum option, int value = 0 )
```

Description: Set an option from the `OgesParameters::OptionEnum` enumerator. See section (3) for a full description of the options available.

option (input) : choose an option

value (input) : value to assign (for options requiring a value).

8.16 set(OptionEnum,float)

```
int  
set( OptionEnum option, float value )
```

Description: Set an option from the OgesParameters::OptionEnum enumerator. See section (3) for a full description of the options available.

option (input) : choose an option

value (input) : value to assign (for options requiring a value).

8.17 set(OptionEnum,double)

```
int  
set( OptionEnum option, double value )
```

Description: Set an option from the OgesParameters::OptionEnum enumerator. See section (3) for a full description of the options available.

option (input) : choose an option

value (input) : value to assign (for options requiring a value).

8.18 set(SolverEnum)

```
int  
set( SolverEnum option )
```

Description: Select a solver from the OgesParameters::SolverEnum enumerator. See section (3) for a full description of the options available.

option (input) : option selected.

8.19 set(SolverMethodEnum)

```
int  
set( SolverMethodEnum option )
```

Description: Select a solver method from the OgesParameters::SolverMethodEnum enumerator. See section (3) for a full description of the options available.

option (input) : option selected.

8.20 set(PreconditionerEnum)

```
int  
set( PreconditionerEnum option )
```

Description: Select a preconditioner from the OgesParameters::PreconditionerEnum enumerator. See section (3) for a full description of the options available.

option (input) : option selected.

8.21 set(MatrixOrderingEnum)

```
int  
set( MatrixOrderingEnum option )
```

Description: Select a matrix ordering from the OgesParameters::MatrixOrderingEnum enumerator. See section (3) for a full description of the options available.

option (input) : option selected.

8.22 get(OptionEnum,int&)

```
int
get( OptionEnum option, int & value ) const
```

Description: Return the current value of an option (this version appropriate for options that have a value of type ‘int’. See section (3) for a full description of the options available.

8.23 get(OptionEnum,real&)

```
int
get( OptionEnum option, real & value ) const
```

Description: Return the current value of an option (this version appropriate for options that have a value of type ‘real’. See section (3) for a full description of the options available.

8.24 setOgesParameters

```
int
setOgesParameters( const OgesParameters & par )
```

Description: Assign the values from an OgesParameters object to an Oges object.

8.25 sizeOf

```
real
sizeOf( FILE *file =NULL ) const
```

Description: Return number of bytes allocated by Oges; optionally print detailed info to a file

file (input): optionally supply a file to write detailed info to. Choose file=stdout to write to standard output.

Return value: the number of bytes.

8.26 printStatistics

```
int
printStatistics(FILE *file = stdout) const
```

Description: Output any relevant statistics

8.27 updateToMatchGrid

```
int
updateToMatchGrid( CompositeGrid & cg0 )
```

Purpose: Give Oges a new matrix to use. Use this routine, for example, when a grid has moved. This routine will cause the matrix to be refactored the next time solve is called.

cg0 (input): use this CompositeGrid

8.28 updateToMatchGrid

```
int
updateToMatchGrid( MappedGrid & mg )
```

Purpose: Use this version when you are solving a problem on a MappedGrid.

mg (input): use this MappedGrid

8.29 getMatrix

```
int  
getMatrix( IntegerArray & ia_, IntegerArray & ja_, RealArray & a_,  
          SparseStorageFormatEnum format =compressedRow)
```

Description: Return the matrix in a given format.

ia_,ja_,a_ (output) : reference to the matrix in sparse form.

format (input): sparse format

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